Please substitute the following claim set for the pending claim set.

- 1-89. (Cancelled)
- (Currently amended) A method of karyotyping a genome of a test eukaryotic cell, comprising:

identifying pieces of the genome of the test eukaryotic cell by determining nucleotide sequence of said pieces;

enumerating the pieces within a plurality of windows of fixed size of the genome, each of said windows having the fixed size, wherein each window comprises a plurality of pieces and the pieces within a window are genomically clustered;

performing a plurality of comparisons for the plurality of windows in which a first number of pieces enumerated within a window for the test eukaryotic cell is compared to a second number of pieces enumerated within the window for a reference eukaryotic cell, wherein a difference between the first number and the second number indicates a karyotypic difference between the test eukaryotic cell and the reference eukaryotic cell.

- 91. (Canceled)
- (Currently amended) The method of claim 90 wherein the pieces within the each window map within 40 kb.
- (Currently amended) The method of claim 90 wherein the pieces within the each window map within 200 kb.
- (Currently amended) The method of claim 90 wherein the pieces within the each window
 map within 600 kb.

- (Currently amended) The method of claim 90 wherein the pieces within the each window
 map within 4 Mb.
- 96. (Currently amended) The method of claim 90 wherein the piece is pieces are defined by the presence of a BegI restriction endonuclease recognition site which is flanked by 12 nucleotides on either end.
- 97. (Previously Presented) The method of claim 90 further comprising the step of: identifying aneuploidy if (a) pieces of one or more autosomes are determined to be present in the test eukaryotic cell relative to the reference eukaryotic cell at a ratio of 1.5 or greater or less than 0.7; or (b) pieces of one or more sex chromosomes in a male are determined to be present in the test eukaryotic cell relative to a reference male eukaryotic cell at a ratio of 1.5 or greater or less than 0.7; or (c) pieces of X chromosomes in a female are determined to be present in the test eukaryotic cell relative to a reference male eukaryotic cell at a ratio of 3 or greater or less than 1.5 or relative to a reference female eukaryotic cell at a ratio of 1.5 or greater or less than 0.7.
- 98. (Previously Presented) The method of claim 90 wherein pieces representing less than 15 % of the genome of the eukaryotic cell are enumerated in the step of enumerating.
- (Currently amended) A method of determining changes in copy number of portions of the genome of a test eukaryotic cell, comprising:

identifying pieces of the genome of the test eukaryotic cell by determining nucleotide sequence of said pieces; enumerating the pieces within a plurality of windows of fixed size of the genome, each of said windows having the fixed size, wherein each window comprises a plurality of pieces and the pieces within a window are genomically clustered;

performing a plurality of comparisons for the plurality of windows in which a first number of pieces enumerated within a window for the test eukaryotic cell is compared to a second number of pieces enumerated within the window for a reference eukaryotic cell, wherein a difference between the first number and the second number indicates a change in copy number of a portion of the genome between the test eukaryotic cell and the reference eukaryotic cell.

- 100. (Previously Presented) The method of claim 99 wherein the change in copy number is due to gain or loss of a whole chromosome.
- 101. (Previously Presented) The method of claim 99 wherein the change in copy number is due to a gain or loss of a chromosomal arm.
- 102. (Previously Presented) The method of claim 99 wherein the change in copy number is due to an interstitial amplification.
- 103. (Previously Presented) The method of claim 99 wherein the change in copy number is due to an interstitial deletion
- 104. (Currently amended) The method of claim 99 wherein the pieces within the each window map within 40 kb.
- 105. (Currently amended) The method of claim 99 wherein the pieces within the each window map within 200 kb.
- 106. (Currently amended) The method of claim 99 wherein the pieces within the each window map within 600 kb.

- 107. (Currently amended) The method of claim 99 wherein the pieces within the each window map within 4 Mb.
- 108. (Currently amended) The method of claim 99 wherein the piece is pieces are defined by the presence of a BcgI restriction endonuclease recognition site which is flanked by 12 nucleotides on either end.
- 109. (Previously Presented) The method of claim 99 further comprising the step of: identifying aneuploidy if (a) pieces of one or more autosomes are determined to be present in the test eukaryotic cell relative to the reference eukaryotic cell at a ratio of 1.5 or greater or less than 0.7; or (b) pieces of one or more sex chromosomes in a male are determined to be present in the test eukaryotic cell relative to a reference male eukaryotic cell at a ratio of 1.5 or greater or less than 0.7; or (c) pieces of X chromosomes in a female are determined to be present in the test eukaryotic cell relative to a reference male eukaryotic cell at a ratio of 3 or greater or less than 1.5 or relative to a reference female eukaryotic cell at a ratio of 1.5 or greater or less than 0.7.
- 110. (Previously Presented) The method of claim 99 wherein pieces representing less than 15 % of the genome of the eukaryotic cell are enumerated in the step of enumerating.